## **AMENDMENT TO THE CLAIMS**

Please amend the claims as follows:

Claims 1-15 (Canceled).

16. (Currently Amended) A method comprising:

obtaining a plurality of images of a measured fluorescence intensity decay for a sample having been exposed to an excitation pulse generated by an excitation light source, the measured fluorescence intensity decay being associated with a fluorescence decay function and/or a fluorescence impulse response function;

deconvolving the excitation pulse from the measured fluorescence intensity delay;

expanding the fluorescence decay function and/or the fluorescence impulse
response function on a Laguerre basis;

estimating a first expansion coefficient (" $\{e_{\theta}\}$ ") of a plurality of expansion coefficients (" $\{e_{j}\}$ ") within the Laguerre basis at each pixel of a plurality of pixels in an the images and computing a map of the first expansion coefficient (" $\{e_{\theta}\}$ ");

generating a map of the higher expansion coefficients cach of the plurality of expansion coefficients (" $\{c_j\}$ "); and

computing a map of <u>average</u> lifetimes by constructing an impulse response function ("IRF") at every pixel for a predetermined number of time instances ("S") and interpolating a time point at which the IRF becomes 1/e of its maximum value, wherein the IRF is represented by the equation:

$$h(r,n) = \sum_{j=0}^{L-1} c_j(r) \cdot b_j^{\alpha}(n), n = 0,1,...,S-1$$

- 17. (Original) The method of claim 16, wherein the sample is selected from the group consisting of a biological tissue, a chemical, a biochemical sample and combinations thereof.
- 18. (Original) The method of claim 16, further including detecting a physiological

condition from the group consisting of a tumor and an atherosclerotic plaque.

- 19. (Original) The method of claim 16, further including predicting the distribution of concentration of at least one biochemical component of the sample images, wherein the sample is composed of a plurality of biochemical components.
- 20. (Original) The method of claim 16, further including monitoring an intracellular component and an activity of the intracellular component.
- 21. (Original) The method of claim 16, further including identifying a chemical with a biological activity for automated screening of the sample for new drugs discovery.
- 22. (Previously Presented) The method of claim 21, further configured to characterize drugs based on their chemical composition so high speed/throughput surveying and counting of the drugs is possible.
- 23. (Previously Presented) The method of claim 21, further configured to characterize a biochemical essay based on biochemical contents to facilitate high speed/throughput surveying/analysis of the essay.
- 24. (Original) The method of claim 16, further including sequencing a deoxyribonucleic acid (DNA) microarray.

Claims 25-44 (Canceled).